

Hepatitis C Treatment Prior Authorization (PA) Request Form

Fax completed form and supporting documentation to 1-800-424-5881

See the Preferred Drug List (PDL) page 11-14	for full Hepatitis C PA o	criteria at: <u>https://w</u>	<u>ww.colorado.go</u>	v/hcpf/pharmacy-resources.
Member name:		DOB:	Medicaid	ID:
Gender: male □ female □ Is the w	voman of childbea	ring potential?	□ No □ Yes	
If yes, have pregnancy test results				ed treatment start
date, and counseling has been pro			-	
-		- ,		-
*If patient is less than 18 years, ir	iulcate patient we	agnic (for dosing	ــــــــــــــــــــــــــــــــــــــ	kg of ibs (circle offe)
Physician:	Phone:	Fax: _		_ NPI:
Prescriber signature (required):			Date:	
Is the prescriber an infectious disease	e specialist, gastroe	nterologist, or h	epatologist?	□ No □ Yes
If no, is the requested drug being pre	escribed by a prima	ry care provider	(PCP) in cons	sultation with an
infectious disease specialist, gastroen	iterologist, or hepat	cologist (CIRCLE	one)? \square N	o □ Yes
If yes, provide first and last name of				
If no, is the requested drug being pre			er without cirr	hosis, by a PCP who has
completed the HCV ECHO series (CIR	CLE one)? 🗆 No	□ Yes		
1- Has member previously received	d direct acting ant	tivirals (DAAs)	or been trea	ted for Hepatitis C?
If yes, fill in Re-treatment box and co	omplete #1-12. If no, fi	ill in Initial treatmen	t box and compl	ete #1-19. No Yes
<u>Pre-treatment</u>				
2- Patient records are attached indi	-		, ,	
(Due to risk of HBV reactivation with DAAs monitor for HBV in all patients receiving D		ionals should scre	en and	□ No □ Yes
3- Physician attests to meeting one of	•	sk opo):		
Member has a diagnosis of chro	5 \	•	JA viral load fo	r > 6 months) □
 Member has a diagnosis of acut 	**			-
Member will be treated upon in				
spontaneous resolution of acute				
therapy (acute HCV infection m	ay spontaneously cle	ar in 20-50% of p	atients) 🗆	
4- Provider attests that member is	ready to be comp	liant to the me	dication regi	imen □ No □ Yes
 Prescribers should utilize assessn 	nent tools to evaluate	readiness of the	patient for trea	atment
 https://www.thenationalcouncil 		ds/2020/04/Screen	ning-for-Viral-He	patitis-within-Behavioral-
Health-Organizations-7.9.14.pdf	·			
Psychosocial Readiness Evaluation				
5-Member's complete current medicati	-		_	
interactions have been screened for ar	•	fore and during	treatment.	□ No □ Yes
May use https://www.hep-drugint			_	
6- Is the member is abusing/misusing				□ No □ Yes
6a-If yes, does the provider attest that				
use treatment program, prior to initia	_		□ No □ Yes	
Treatment referrals can be reque Treatment referrals can be reque				
calling customer service accessib organizations	ic at. <u>iittps://www.ne</u>	zaiu ii ii Stcoiorado.(COMPREADED - ITS	oc-coloi au0-1 Egi0fidi-
<u>organizaciono</u>				



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- 7- Required lab tests (taken within past 6 mo) are submitted with this request:
 - Quantitative HCV RNA viral load
 - Complete Blood Count (CBC)
 - Hepatic Function Panel (i.e. albumin, total and direct bilirubin, alanine aminotransferase (ALT), aspartate aminotransferase (AST), and alkaline phosphatase levels)
 - Calculated glomerular filtration rate (GFR)
 - If cirrhosis is present, calculation of the Child-Turcotte-Pugh (CTP) Score
 - Transplant status as applicable (pre-, post-, N/A)
- 8- Liver fibrosis test is submitted (this is not required, but, if available): \square No \square Yes Post Treatment:
- 9-Provider attests to provide one HCV RNA test result from 12-24 weeks post-treatment $\ \square$ No $\ \square$ Yes
- Please submit <u>Health First Colorado HCV Treatment Outcomes Form</u> (accessible from the Pharmacy Resources Page)

Initial treatment requests (Fill in requested drug regimen and duration in table below)

Drug	Stren gth/ Formu lation *	Duration (weeks)	Preferred Initial Treatment Regimens (GT-Genotype, NC-No-Cirrhosis, CC-Compensated Cirrhosis, DC-Decompensated Cirrhosis)
ledipasvir/ sofosbuvir (Harvoni)			Members 3 years and older for GT 1, 4-6 with NC or CC; or GT 1 in combination with ribavirin in DC; or GT 1,4 in combination with ribavirin for liver transplant recipients with NC or CC. If request is for pellets, member is 3 years of age or older weighing less than 17kg OR 3 years or older that are unable to take/swallow ledipasvir/sofosbuvir oral tablets
Mavyret	100mg -40mg		Members 12 years and older or weighing at least 45 kg with NC or CC (Child-Pugh A only)
Sofosbuvir/ velpatasvir (Epclusa)			Members 6 years and older or weighing at least 17 kg for with NC or CC (Child-Pugh A only); or in combination with ribavirin in DC

Retreatment or prior exposure to DAAs (Fill in requested drug regimen and duration in table below)

Drug	Strength/	Duration	Preferred Regimens For Retreatment or treatment experienced
	Formulation*	(weeks)	(GT-Genotype, NC-No-Cirrhosis, CC-Compensated Cirrhosis, DC-Decompensated
			Cirrhosis)
Mavyret	100mg-40mg		Members 12 years and older or weighing at least 45 kg with NC or CC (Child-
			Pugh A only); or for GT 1, who previously have been treated with a regimen
			containing an HCV NS5A inhibitor or an NS3/4A protease inhibitor (PI), but not
			both
Sofosbuvir/			Members 6 years and older or weighing at least 17 kg with NC or CC (Child-
velpatasvir (Epclusa)			Pugh A only); or in combination with ribavirin in DC
Vosevi	400mg-		Members 18 years or older with chronic HCV infection with NC or CC (Child-
	100mg-100mg		Pugh A only) and either previously failed treatment with a regimen containing
			an NS5A inhibitor (such as ledipasvir, daclatasvir, or ombitasvir)
			OR are GT 1a or 3 and previously failed treatment with a regimen containing
			sofosbuvir without an NS5A inhibitor

10- List previous tre	eatment regime	en received	, and (date received:	·		
11- Genotype of firs	st treatment an	d current g	enoty	pe (if known)	Previous	Current	
12- Was the entire	treatment regir	nen comple	eted?	□ No □ Yes	If no (early	discontinuation	occurred)
olease describe:							

- Adverse effects experienced from previous treatment regimen
- Concomitant therapies during previous treatment regimen



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3rd page only needs to be attached if applicable to request.

Ν	lo	n-	Pr	efe	rre	be	D	AAs
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If not prescribing a preferred treatment regimen, provide rationale and supporting documentation (Acceptable rationale may include member has initiated treatment on a non-preferred drug and needs to complete therapy, patient-specific medical contraindications to a preferred treatment).					
Ribavirin (Note: Preferred ribavirin products do not require a PA)					
Is member ineligible for ribavirin? No Yes					
If so, please provide documentation and medical notes for consideration of approval.					

Does the requested regimen include ribavirin? □ No □ Yes

If yes, Provider Attests to the following:

- Member is not a pregnant female or a male with a pregnant female partner
- Women of childbearing potential and their male partners must attest that they will use two forms of effective (non-hormonal) contraception during treatment
- Member does not meet any of the following ineligibility criteria for use of ribavirin:
 - Pregnant women and men whose female partners are pregnant
 - Known hypersensitivity to ribavirin
 - Autoimmune hepatitis
 - Hemoglobinopathies
 - Creatinine Clearance < 50mL/min
 - o Co-administered with didanosine